JUN 0 5 2002

Docket No. MCP 264

Applicants

Codispoti, Joseph R.

Serial No.

09/709,069

Art Unit: 1614

Filed

9 November 2000

Examiner: Jagoe, D.

For

EN CONTRACTOR OF THE PARTY OF T METHOD FOR TREATING MIGRAINE SYMPTOMS WITH IBUPROFEN

HE UNITED STATES PATENT AND TRADEMARK OFFICE

I hereby certify that this correspondence is being deposited with the United States Postal Service with sufficient postage as first class mail in an envelope addressed to: Assistant Commissioner for Patents, Washington, DC 20231 on

(Date of Deposit)

Michele G. Mangini (Name of applicant, assignes, or Registered Representative

Assistant Commissioner For Patents Washington, D.C. 20231

SUPPLEMENTAL DECLARATION UNDER 37 CFR 1.131

Dear Sir.

- 1. This Supplemental Declaration is submitted to supplement the Declaration Under 37 CFR 1.131 malled on 25 August 2000 in response to the 30 March 2000 Office Action in the parent application, United During the prosecution of the above-referenced States Serial No. 09/449,124 (herein "Declaration"). application, I became aware of the fact that the Declaration contained an inadvertent typographical error in the page number listed for the Furey Abstract. This inadvertent error is corrected in Paragraph 2 herein.
- 2. This Supplemental Declaration is submitted to establish completion and reduction to practice of the invention in the above-identified application in the United States at a date prior to 24 August 1999. It is my information and belief that the Information Center of McNeil Consumer & Specialty Pharmaceuticals Division of McNeil-PPC, Inc., the assignee of record to the entire right, title, and interest in the aboveidentified application (hereinafter "Assignee"), received a copy of the abstract entitled "Efficacy and Safety of Ibuprofen (I) Liquigels in Migraine Headache: A Randomized, Double-Blind Placebo-Controlled Study by Furey, et al., as published in Volume 39(9) of the Journal of Clinical Pharmacology on page 978 (Sept. 1999) (hereinafter "Furey Abstract"), on or about 24 August 1999. It is further my information and belief that this volume of the Journal of Clinical Pharmacology was mailed to its subscribers on or about 20 August 1999. A copy of the Furey Abstract is attached hereto as Exhibit A. The Furey Abstract was cited in the Office Action mailed on 27 February 2002 in the above-referenced application.

- 3. I, Joseph R. Codispoti, MD, am the sole inventor on the invention described and claimed in the above-identified application.
- 4. As of approximately August 2001 until the present, I am employed by Sanofi-Synthelabo Research and Development located at 9 Great Valley Parkway, Malvern, PA 19355. Previous to that date, and at and before the completion of the invention. I was in the employ of the Assignee.
- 5. I understand that the claims of the present invention have been rejected in view of the Furey **Abstract**
- 6. Appended hereto as Exhibit B is a true copy of the Clinical Study Report entitled "A Single Dose, Randomized, Double-Blind, Placebo-Controlled Study Evaluating the Safety and Efficacy of Ibuprofen 200 mg and 400 mg for the Treatment of Migraine Headache Pain (hereinafter "Report"), which was performed at my request and which memorializes the conception and reduction to practice of the claimed invention.
- 7. On page 12 of the Report, it can be seen that the invention of this application, i.e. a method for mitigation or treating photophobia and phonophobia associated with migraines by providing an effective amount of ibuprofen as the sole anti-migraine agent, was made prior to August, 1999, which is earlier than the 35 USC §102(f) date of the Furey Abstract.
 - 8. All dates that have been redacted in the Exhibit are before August, 1999.

8. I, Joseph R. Codispoti, further declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further declare that the statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under 35 USC §1001, Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application or patent issuing thereon.

Joseph R. Codispoti, MD

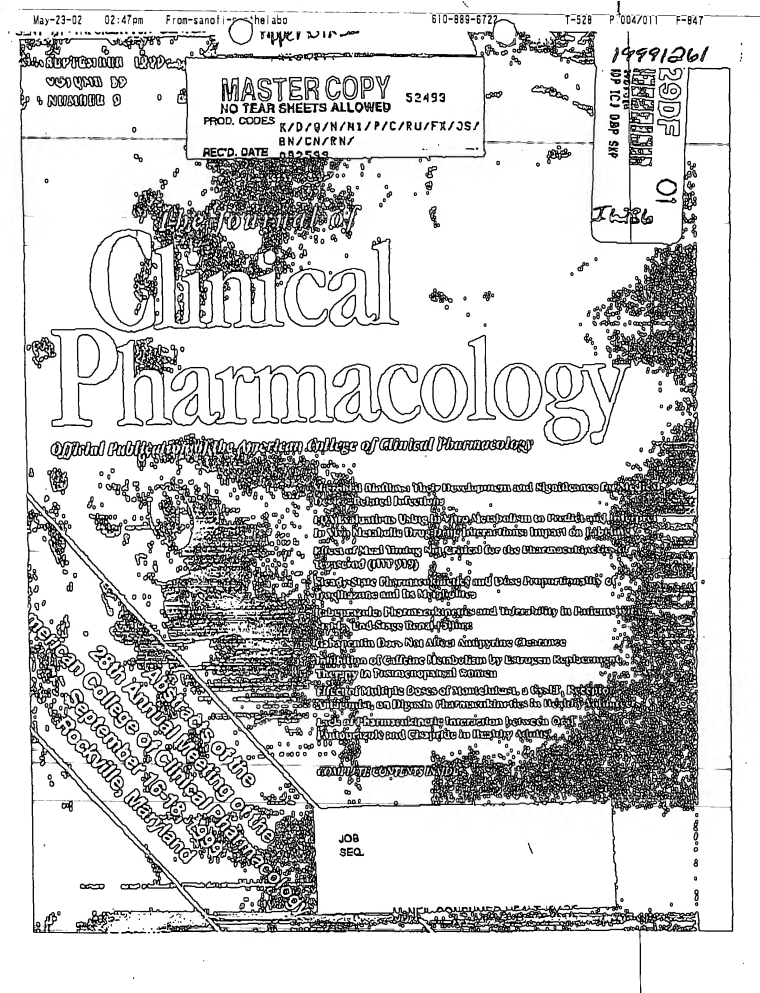
Country of Citizenship: USA

Touthington Rd, Phila, PA

Appendix A: Furey Abstract

Appendix B: Clinical Study Report

Mcp264-131decn.doc



ExhibitA

TWENTY-BIGHTH ANNUAL ACCP MEETING ABSTRACTS

34

EFFICACY AND SAFETY OF BUPECIPEN (I) LIQUIDELS IN MCBAINE HEADACHE: A RANDONDZED, DOUBLE-BLIND, PLACEBO-CONTROLLED STUDY. S.A. FROM! II Kellicia" », R. Gathi", B. Ar", P. Consillo" «I. Toper" ». "Modical Department, Whithyll-Bohins Healthout, Medical, NJ 2nd "Michigan Head-Priza & Naurological Legislots, Ann Arbor, MI.

We compared I string and I coored ministered as impaged to placed (PBO) among subjects with moderate of avere migrate medical with moderate of avere migrates medical arbitrar among and microsity on a four-paid and roled pain after define public and roled pain microsity on a four-paid and roled pain relict on a five-point categories tools. Subjects also meed the associated uniquales symptoms of among almostophobia, and arsessed their limitation of activity (LOA) using a quality of life index singless from 0 (among to 3 (avero). The promoter of the control of a control of the control of

Pa dpa)ne	PED Photog	(40700)	(Street)
Cumulative respondent 2h (%)	47	270	390
SPRID (GIGN)	7.0	0,90	30.70
Medica) a perceptitio relet (min)	66	470	6Q*
LOA IMPORTATE (6h)	0.0	3,02	Q Da
Namena Improvement (0b)	0.2	0.3	0,5=

(* peolit 72 fBO); I goon; and coons was both algolitemity aperior to PBO in the emissions. We adopted this so assure, placeptatia, and indusephobia over the I George tended to appoint a multiplicated with adverse experience furthered comparator of FBO: Rised on these data, we expelied that I depoint affects of FBO: Rised on these data, we expelied that I depoint of the property edictors allowing effectives and emproves allowing the number ascribing angular symptoms, and improves migratican, facility of life, the abulton descriptions and improves migratican, facility of life, the abulton of the contemporary and the profession allowing done.

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PHARMACOKINETICS OF FMD 112 13R AFTER ESCALATING SINGLE ORAL DOSES OF GANTOPIDAN, A NEW GLYCOPROTEIN LIBRILIA RECEPTOR ANTAGOMIST. Bend Melibohm. Roland Neugeboods, Rail-Ulrida Blutings, Michael Schules, and Andrew Kovas. Colloge of Photmacy. University of South Citolina, Columbia, SC. and Clinical Photmacology, Merci KO2A. Darmacod, Geomany.

Cambaldon is an early available cauble product. Dissertivation results in the active metabolic EMD 132 338, a putent, reversible, non-peptide entagonist of the glycoproduct limits acceptor (GPR) for the inhibition of planets aggregation associated with thromboembolic events. In a phase I clinical study, the pharmaenkineries of EMD 132 338 were evaluated in exquential groups of healthy mais subjects after single oral doors of 2.3 (prs), 1.3 (prs),

978 • J Clin Pharmacol 1999;39:969-985

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EFFECT OF NEVERAPINE ON HUMAN BLOOD CLUTATHIONE LEVELS AND SUBCHROPIC TOXICITY AFTER DERMAL ADMINISTRA-TION TO RATS. Chukwueineka S. Okereke. Univ. of Rhode Island College of Pharmacy, Roger Williams Med. Cent. Providence, Bl.

Nevirapine (NVP) is a potent non-nuclediside reverse transcriptuse imbibitor that has been shown to inscriver the human immuno-deficiency virus upon administration. Currently, attempts at reducing incidences of vertical transmission of the virus (undeber to child have focused on the use of phurmacological agents in "birth canal cleansing" during child labor and delivery. Pollowing subchronic administration of HVP to semale this twice dully for 4-works, body weight, clinical chemistry and bematological parameters were not affected. However, invivo blood glumphions (CSH) was reduced Similarly, invitro blood OSH time column in bossers and rate were reduced latitudy up maril 45 minutes and gradually returned to control levels thereafter. The robound in GSH levels is probably due to a compensatory mechanism due to OSH-reductore enzyme. Based on these studies, NVP does not seem to produce any appreciable dermal effects in rats.

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GABAPENTIN SINGLE-DOSE PHARMACOKINETICS IN HEALTRY INFANTS AND CHILDREN GENER M. Hall?, Deward N. Booktroder, Dovid L. Westie*, Sumuel W. Boelber?, Richard Brown?, Noncy Junioezic-Dolphin*, and Edward L. Posver, Purio-Davis Pharmana-affeol Research; Ann Arbor, ML

Gabapentin (Neuronda) is a gamma-amataoburario acid analogue indicated to adults for adjunctive accessors of porial science with or without secondary generalization. Two ejudica ware conducted to determine the tangle-date pharmscoldectico of gatapartia in healthy subjects age I mouth to 12 years and to gaida dona salaction in salaty and efficacy trials to pediatric policate for the above indication. Fore-eight subjects wele given a single close of gobspeciain 10 moving administrated brally while fisting. Emoliment was bomogeneously distributed throughout the age range. Plaume samples were drawn pro-dure, then serially for 24 hours. A single done of goldsports was well colerned by healthy pediatric subjects. Plats of age vs. AUC(0-ब्द) काह्यस्थापने केंग्रीस्थातकस्था in younges (1 month to 4 years) þr. alder (5 to 12 years) subjects. Beans A.VC(0-c) was 25.7 ug ha/ml in younger subjects and 36.0 ug ha/ml in older applients (p<0.001). Classinas (nemesticad as weight) with 7.35 ml/min/kg for younger subjects and 4.41 ml/min/kg for pletr subjects (p<0.001). Merun peak plasma concernations (Crean) were 3.76 and 4.52 racgimi, respectively (p-3.05). Districtions in the colculated biografishility could not sufficiently explicit the dispurity in AUC. Patterns between 1 month and 4 years would require an approximate 30% larger daily dose to achieve direller drog exposures to these pasients between 5 and 12 years of age.

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AppendixII.

MCNEIL CONSUMER HEALTHCARE

CLINICAL STUDY REPORT



PHASE III

A Single-Dose, Randomized, Double-Blind, Placebo-Controlled Study Evaluating the Safety and Efficacy of Ibuprofen 200 mg and 400 mg for the Treatment of Migraine Headache Pain.

START DATE: END DATE: 237 Report Date Report No.

Blostatistician, Date

Statistical Services

Brenda Zimmerman, MS

Assistant Director, Statistical Services

Vanessa Burczynski, BS

Medical Program Administrator,

Clinical Devel pment

Date

James B. Nick, Director,

Codispotk MD

Clini al D vei pment

Exhibit B

Date

Date

Date

610-889-6727

T-528 P.007/011

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Clinical Study Report Ibuproten Tablet 200mg McNeil Consumer Healthcare

1. SYNOPSIS

Name of Sponsor/Company McNeil Consumer Healthcare	Individual Referring Dossier	Study to Park	Table Of the	.(For National Authority Use Onty)
Name of Finished Product: Motin Migraine (Ibuprofen Tablet 200 mg)	Yoluma:	<i>آ</i> وج		·
Name of Active Ingredient:	Paga:			

Title of study: A Single-Dose, Randomized, Double-Blind, Placebo-Controlled Study Evaluating the Salety and Efficacy of Ibuprofen 200 mg and 400 mg for the Treatment of Migraine Headache Pain.

Investigators: The 16 investigators are listed in Section 4, Investigators and Study Administrative Structure

Study Conters: The 18 Investigative sites are listed in Section: 4, Investigators and Study Administrativo Structure

Study Reciaco

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Phase of development: W

Objectives: The purpose of this study was to evaluate the efficacy and safety of fourrolen 200 mg and buprolen 400 mg for the treatment of pain associated with migrains headachs.

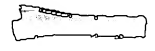
Methodology: This was a multicenter, single-dose, randomized, double-blind, paralici, placebo-consolica study of approximately 600 subjects, 18 years of age and older, experiencing at least moderate pain accordated with migraine headache. Following a screening visit, eligible subjects wars removinly sculptured to elikar louproten 200 mg, ibuproten 400 mg or placebo. Subjects left the investigative center with one dece of bilinded study drug, a timing device, and a subject diary. After the eccumence of a migratine headacho of at least moderate intensity, subjects dosed with study medication and recorded in the diany the date and time of outly medication administration. Efficacy and salety were essessed at specified intervals for six hours following the use of study medication. Subjects returned to the site for a follow-up visit within 72 hours after desired with medication.

Number of subjects:

This study was designed for the completion of at least 600 subjects. Data were available for 649 subjects. Of of whom were included in an intent-to-treat efficacy analysis. All subjects who doesd with study medication and who had officacy data were included in the intent-to-treat analysis. Date were available for 641 subjects in the per-protocol analysis. The table below summarizes the distribution of these subjects by treatment group.

	lbu 200 mg	Ibu 400 mg	Placabo	Total
Enrolled	240	239	23%	719 649
InterT-or-metral	216	219	214 21 3	649
Per-Protocol	214	214		





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Clinical Study Report Ibuprofen Tablot 200mg McNeil Consumer Healthcare

Name of Sponsor/Company McNeil Consumer Healthcare	Individual Referring Dossier	Study to Part	Table of the	(For National Authority Use Only)	
Name of Finished Product: Motrin Migraine (Ibuprofen Tablet 200 mg)	Volume:				
, i	Page:				
Name of Active Ingradient:					
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1					

The table below summarizes the demographic characteristics for all subjects enrolled:

Characteristic	- Ibu 200 mg (N = 240)	Ibu 400 mg (N = 239)	Placebo (N =234)	Total (N = 713)
Sex (n,%) Male Fomale	42 (17.5) 198 (82.5)	35 (14.8) 204 (86.4)	34 (14.5) 200 (85.5)	111 (15.6) 602 (84.4)
Meau ada (Aus)	38.9	38.5	38.2	38.6
Race (n.%) c-Callestan	74 (89.2) 15 (8.2)	200 (83.7) 18 (7.5) 21(8.8)	208 (88.0) 12 (5.2) 18 (6.8)	620 (87.0) 45 (8.3) 48 (8.7)
Ciher				. •

Diagnosis and main criteria for inclusion: Wigrains headache. Subjects were required to have history of one migrains headache every two months to six migraine headaches per month that were not debilitating or incapacitating.

Test product, dose and mode of administration, balch number: Study dug transment was Motion IB, 200 mg and 400 mg, oral tablet, control number C-778-18.

Duration of treatment: Subjects were meated with a single dose of study drug when they experienced a migraine. Subjects were evaluated for six hours after starting treatment. After desing with study medication. subjects returned to the investigative site for a follow-up visit.

Reference therapy, dose and mode of administration, batch number: Reference therapy consisted of an oral placebo tablet, control number C-220-GA.

Efficacy: The primary efficacy endpoint was the percentage of subjects who experienced a reduction in baseline pain Intensity from severe (3) or moderate (2) to mild (1) or none (0) at the two hour postmedicarden assessment time (defined as 'responders'). An additional primary efficacy endpoint was the pain imenally difference from baseline at two hours. Secondary measures of efficacy included: percentage of subjects pain fine at two hours; percentage of subjects with associated migrains symptoms reduced to zero at two and elehours; time to rescue and reacue rate; pain intensity differences from baseline and pain relief from 0.5 to 8 hours. SPID, TOTPAR, severity differences from baseline for the associated migrains symptoms from 0.5 to 6 hours; emergence of associated symptoms; subject rating of overall impression of medication; and time to and intensity of recurrent headaches.

Safety: Safety assessments consisted of a routine physical examination at baseline and monitoring of adverse events.

Clinical Study Report Ibuprolen Tablet 200mg McNell Consumer Healthcare



ĺ	Name of Spons r/Company	Individual	Study		(For National	
	McNeil Consumer Healthcare	Referring	to Part	of the	Authority Use Only)	l
ì		Dossier				ļ
	Name of Finished Product:					١
	Motrin Migraine	Volume:				h
	(Ibuprofen Tablet 200 mg)					II.
		Page:				h
	Name of Active Ingredient:	_				I
	ibuprofen					N
]	V

Statistical Methods: There were three pairwise comparisons of interest for analysis: ibuprolen 200 mg vs. placebo, ibuprolen 400 mg. Each of the statistical tests described below were performed for each treatment pair at the 0.05. two-tall alpha level. The Intent-to-treet analysis was the primary analysis.

Primary Measures:

A Cochran-Mantel-Maenszel test of general association stratified by baseline tevel of pain intensity was used to make painwise treatment comparisons of response rates. A three-way ANOVA (Treatment, Baseline Pain, Investigator) was used in the analysis of pain intensity difference (PID) from baseline at two hours; painwise treatment comparisons were made using Fisher's protected LSD technique.

Additional pain measures:

The percent of subjects who were pain free was analyzed with a Cochran-Mantel-Hagnszel test of general association, stratified by initial level of pain intensity. PIDs at times other than two hours and SPID were analyzed similarly to the analysis of PID at two hours. A two-way ANOVA (Treatment, Investigator) was used for the analysis of pain relief (PR) at each time point; TOTPAR was analyzed similarly.

Associated symptoms:

For subjects reporting each symptom at baseline, differences from baseline in sevarity of nauseal photophobia, photophobia, and functional disability at each measurement interval during the six-hour follow-up particle were analyzed using analysis techniques identical to those outlined for PID above with the exception that the beseline severity of each individual symptom was included in the ANOVA model in place of baseline headeache pain intensity. The rates of emergence of each associated symptom after baseline were analyzed using Fish 1's exact tests. Paintine treatment comparisons of the percentage of subjects with the severity of ratesea, photophobia, phonophobia, and functional disability reduced to "none" at two and alt hours were analyzed with Cochran-Mantel-Heenszel tests of general association stratified by baseline level of each symptom. The incidence of vomiting combined across all measurement intervals was compared using Fisher's Exact tests.

Other measures:

Pairwise treatment comparisons for the overall impression of the study medication were made using the extended Cochran-Mantel-Haenszel test with mean modified right scores, stratified by initial lavel of pain intensity. Pairwise treatment comparisons of time to recurrence of migraine headache were performed using the Wissian test available in the SASO LIFETEST procedure. Only subjects who were "respondent" at two hours and had a recurrence of moderate or severe migraine were included in the analysis. Pairwise treatment comparisons of severity of the pain associated with the recurrent migraine headache were analyzed using a Cochron-Mantel-Haenszel test of general association, stratified by initial level of pain intensity. Only subjects with a recurrent migraine headache were included in this enalysis.

Pairwise differences in the survival distributions between treatments for the time to recess were conducted using the Wilcown test available in the SAS° LIFETEST procedure. Rescue rates at six hours were enalyzed using a Cochtan-Mantel-Haenszel test, stratified by initial level of pain intensity.

Subgroup analyses:

The two primary measures were analyzed by baseline pain, gender, and race. In addition, the percentage of responders at two hours was analyzed by mensional status (yes/no).

Salety Modaring:

The frequency of adverse events and frequency of withdrawal from the study were compared between freetment groups with chi-square tests.

610-889-6727

Name of Sponsor/Company
McNeil Consumer Healthcare

Name of Finished Product:
Motion Migraine
(Ibuprofen Tablet 200 mg)

Name of Active Ingredient:
ibuprofen

Efficacy Results: Key demographic and baseline characteristics of the intent-to-treat population are given below:

Characteristic	lbu 200 mg (N = 216)	lbu 400 mg (N ⇒ 219)	Placebo (N = 214)	Total (N = 649)
Sex (n,%) Male Female	36 (16.7) 180 (83.3)	33 (15.1) 186 (84.9)	29 (13.6) 185 (86.4)	98 (15.1) 551 (84.9)
Mean Age (yrs)	38.8	38.5	38.5	38.6
Race (n,%) White African-American Other	191 (88.4) 14 (6.5) 11 (5.1)	185 (84.5) 15 (6.8) 19 (8.7)	191 (89 <i>.</i> 2) 11 (5.1) 12 (5.6)	567 (87.4) 40 (8.2) 42 (6.5)
Baseline Paln (n,%) Moderate Severe	144 (66.7) 72 (33.3)	158 (72.1) 61 (27.9)	152 (71.0) 62 (29.0)	454 (70.0) 195 (30.0)

The key efficacy results from this study are summarized in the table below:

THE KDY CHASSES TO SEE					Significano	6 2
	lbu	lbu 400	Placebo	150 200 Va P186850	1bu 400 vs Placabo	10u 200 V3 10u 400
Endpoint	200	41.10	26.60	S	3	198
Pain to mild or none of 2 hours" (%)	39.81			s	S	NS
Baseline Pain = Moderate	49.31	46.57	28.95	_	N/B	ar
	20.89	29 <u>.</u> \$1	20.57	ris 8	•	M8
Baseline Paln = Severe	6.67	38.6	0.35	3	3	
one of a mean)			0.14	s	Ş	MS
Baseline Pain = Moderate	0.58	0.51			NS	NS.
	0.91	1.02	88.0	R S		NS
Baseline Pain = Severe	13.43	15.98	8.54	S	S	
Pain to none at 2 hours (%)			2.05	6	\$	NS
SPID (mean)	4.17	4.01	-	_	s	RIS.
	9.63	9.52	6.65	\$	_	NS
TOTPAR (mean)	1.14	1.14	0.66	S	S	
Overall Impression of Medication (mean)		•		NS	NS	КS
Явсителсе within 24 hours (%)	31.4	31.1	33.3			

a: S: p < 0.05; NS: p > 0.05.

b: Primary endoord

610-889-672 T-528 P-0117011

Clinical Study Report Ibuproten Tablet 200mg McNoil Consumer Healthcare

(For National Table Study Individual Name of Sponsor/Company Authority Use Only) Referring to Part of the McNeil Consumer Healthcare Dossier Name of Finished Product: Volume: Motion Migraine (Ibuprofen Tablet 200 mg) Page: Name of Active Ingredient: ibuprofen

In addition to these results, there was a significantly greater reduction from baseline in mean severity of migraine-associated symptoms of photophobia and functional disability in both ibupraten groups compared t placebo at all time points in the interval from two to six hours after dosing. For phonophobia, mean severity differences were significant only for the 400 mg ibupraten dose relative to placebo from one to six hours and for nausea, there were no differences between treatments at any time interval.

Safety Results: Ibuproten was well tolerated and no safety issues were identified in this migraine headache population. Overall 34.8% of subjects reported adverse events; there was no significant difference among treatment groups. In addition, drug-related adverse events were reported by 24.7% of study subjects; there was no significant difference among treatment groups. The most common adverse events were in the digestive system (mainly nauses and vomiting), occurring in 30.2% of study subjects. There was no significant difference among treatment groups; it is interested most likely that these symptoms represent the normal sequelae of a migraine headache attack. No serious adverse events or deaths were reported. Three subjects discontinued therapy due to adverse events, two subjects in the ibuprofen 400 mg group and one subject in the placebo group.

Conclusions: Ibuprolen at OTC doses of 200 mg and 400 mg is an effective treatment for the temporary relief of migraine headache pain and the associated symptoms of migraine including photophobia and functional disability.

Efficacy results for subjects with severe migraine pain intensity are not inconsistent with the current labeling regarding OTC louprolen dosing which directs consumers to take 400 mg if pain does not respond to 200 mg.

All secondary efficacy measures including pain reliet and pain intensity difference showed effects consistent ক্ষাঁণ the primary efficacy outcome measures.

lbuproten was well tolerated and no safety issues were identified in this migraine headache population. There were no algorificant differences between either dose of ibuproten and placebo in the incidence of adverse events. The seventy and nature of adverse events were similar among groups. No serious adverse events or deaths were reported.

Date of the report: